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⑯ Heat shrinkable films containing single site catalyzed copolymers having long chain branching.

⑯ The present invention is directed to a heat-shrinkable, thermoplastic film or bag which contains at least one homogeneous ethylene alpha-olefin copolymer with at least some limited long chain branching. Oriented films made in accordance with the present invention exhibit improved processability as well as improved physical properties such as excellent optics and impact resistance. A homogeneous ethylene alpha-olefin with long chain branching may be present in a monolayer film either alone or in a blend or may be included in one or more layers of a multilayer film in accordance with the present invention.

EP 0 600 425 A1

5 Field of the Invention

The present invention is generally directed to heat shrinkable films containing single site catalyzed copolymers which include at least some long chain branching.

10 Background of the Invention

Shrinkable thermoplastic films have found many useful applications in packaging of meats, cheeses, poultry and numerous other food and non-food products. There is always the search for improvement in these films to make them have better impact resistance, improved optics and improved shrinkability. For 15 example, in U.S. Patent No. 4,640,856 to Ferguson et al., the multilayer thermoplastic heat shrinkable film was described having improved shrink, toughness and barrier properties. The film included at least one layer of a very low density polyethylene and a gas barrier layer of vinylidene chloride copolymer or ethylene vinyl alcohol. The film was found to be particularly useful for making bags for packaging large cuts of fresh red meat.

20 U.S. Patent Nos. 5,059,481, 4,976,898 and 4,863,769, all to Lustig et al., disclose heat shrinkable film suitable for packaging food articles such as frozen poultry, primal meat cuts and processed meat products wherein the film may be a biaxially stretched monolayer film of a very low density polyethylene copolymer or a multilayer film containing very low density polyethylene.

25 U.S. Patent No. 4,457,960 to Newsome discloses the use of linear low density polyethylene in multiple layer molecularly oriented films.

Each of the foregoing patents describe the incorporation into heat shrinkable films of conventional ethylene/alpha-olefins produced by Ziegler-Natta catalyst systems. Ziegler-Natta catalytic methods are commonly used throughout the polymer industry and have a long history tracing back to about 1957.

30 These systems are often referred to as heterogeneous since they are composed of many types of catalytic species each at different metal oxidation states and different coordination environments with ligands. Examples of Ziegler-Natta heterogeneous systems include metal halides activated by an organometallic co-catalyst, such as titanium or magnesium chlorides complexed to trialkyl aluminum and may be found in patents such as U.S. Patents 4,302,565 and 4,302,566. Because these systems contain more than one catalytic species, they possess polymerization sites with different activities and varying abilities to 35 incorporate comonomer into a polymer chain.

35 The result of such multi-site chemistry is a product with poor control of the polymer chain architecture both within the sequence of a single chain, as well as when compared to a neighboring chain. In addition, differences in catalyst efficiency produce high molecular weight polymer at some sites and low molecular weight at others. Therefore, copolymers produced using these systems lead to polymer products which are 40 mixtures of chains some high in comonomer and others with almost none. For example, conventional Ziegler-Natta multi-site catalysts may yield a linear ethylene/alpha-olefin copolymer (HDPE, LLDPE, VLDPE, ULDPE) having a mean comonomer percentage of 10, but with a range of 0% to 40% comonomer in individual chains. This, together with the diversity of chain lengths results in a truly heterogeneous mixture also having a broad molecular weight distribution (MWD).

45 Linear low density polyethylene (LLDPE) has enjoyed great success as a raw material choice for packaging films. The term LLDPE is generally understood to describe copolymers of ethylene and one or more other alpha olefin monomers which are polymerized at low pressure using a Ziegler-Natta catalyst to achieve a density range of about 0.915 to about 0.940. Although no clear standard exists, LLDPE polymers are often marketed in subgroups of densities such as linear medium density (LMDPE), linear low density 50 polyethylene, linear very low density (VLDPE), or linear ultra low density polyethylene (ULDPE). These classifications are for marketing use and will vary by supplier.

55 These materials are different from high pressure low density polyethylene (LDPE) which is generally understood in the trade as a highly branched homopolymer having a single low melting point. For example, a 0.92 density LDPE would typically have a melting point at about 112°C while a corresponding density LLDPE would have melting points at 107°, 120°, and 125°C. The multiple melting points are commonly observed with LLDPE and are a consequence of the above mentioned heterogeneous incorporation of comonomer.

Recently a new type of ethylene copolymer has been introduced which is the result of a new catalyst technology. Examples of introductory journal articles include "Exxon Cites 'Breakthrough' in Olefins Polymerization," Modern Plastics, July 1991, p.61; "Polyolefins Gain Higher Performance from New Catalyst Technologies," Modern Plastics, Oct. 1991, p.46; "PW Technology Watch," Plastics World, Nov. 1991, p. 29; and "Plastics Technology, Nov. 1991, p. 15.

Such new resins are produced using metallocene catalyst systems, the uniqueness of which resides in the steric and electronic equivalence of each catalyst position. Metallocene catalysts are characterized as having a single, stable chemical type rather than a volatile mixture of states as discussed for conventional Ziegler-Natta. This results in a system composed of catalyst positions which have a singular activity and selectivity. For this reason, metallocene catalyst systems are often referred to as "single site" owing to the homogeneous nature of them, and polymers and copolymers produced from them are often referred to as single site resins by their suppliers.

Generally speaking, metallocene catalysts are organometallic compounds containing one or more cyclopentadienyl ligands attached to metals such as hafnium, titanium, vanadium, or zirconium. A co-catalyst, such as but not limited to, digomeric methyl alumoxane is often used to promote the catalytic activity. By varying the metal component and the cyclopentadienyl ligand a diversity of polymer products may be tailored having molecular weights ranging from about 200 to greater than 1,000,000 and molecular weight distributions from 1.5 to about 15. The choice of co-catalyst influences the efficiency and thus the production rate, yield, and cost. Examples of metallocene catalysts are disclosed in U.S. Patent Nos. 4,701,432, 4,306,041, 5,088,228, 4,935,397, 5,084,534, 3,161,629, 5,055,438, 5,057,475, and in JP 63/175004 and JP 1,101,315.

As a consequence of the single site system afforded by metallocenes, ethylene/alpha-olefin copolymer resins can be produced with each polymer chain having virtually the same architecture. Therefore, the copolymer chains produced from single site systems are uniform not only in chain length, but also in average comonomer content, and even regularity of comonomer spacing, or incorporation along the chain.

In contrast to the above mentioned Ziegler-Natta polymers, these single site metallocene polymers are characterized as having a narrow MWD and narrow compositional distribution (CD). While conventional polymers have MWD's of about 3.5 to 8.0, metallocenes range in MWD from about 1.5 to about 2.5 and most typically about 2.0. MWD refers to the breadth of the distribution of molecular weights of the polymer chains, and is a value which is obtained by dividing the number-average molecular weight into the weight-average molecular weight. The low CD, or regularity of side branches chains along a single chain and its parity in the distribution and length of all other chains, greatly reduces the low MW and high MW "tails". These features reduce the extractables which arise from poor LMW control as well as improve the optics by removing the linear, ethylene-rich portions which are present in conventional heterogeneous resins.

Thus, conventional Ziegler-Natta systems produce heterogeneous resins which reflect the differential character of their multiple catalyst sites while metallocene systems yield homogeneous resins which, in turn, reflect the character of their single catalytic site.

Another distinguishing property of single site catalyzed ethylene copolymers is manifested in their melting point range. The narrow CD of metallocenes produces a narrow melting point range as well as a lower Differential Scanning Calorimeter (DSC) peak melting point peak. Unlike conventional resins which retain a high melting point over a wide density range, metallocene resin melting point is directly related to density. For example, an ethylene/butene copolymer having a density of 0.905 g/cc produced using a metallocene catalyst has a peak melting point of about 100°C, while a slightly lower density ethylene/butene copolymer which was made using a conventional Ziegler catalyst reflects its heterogeneous nature with a melting point at about 120°C. DSC shows that the Ziegler resin is associated with a much wider melting point range and actually melts higher despite its lower density.

It should be noted that at least some previously available ethylene based linear polymers approximated the physical and compositional properties achieved by the present metallocene catalyzed polyolefins. For example, in "Sequence and Branching Distribution of Ethylene/1-Butene Copolymers Prepared with a Soluble Vanadium Based Ziegler-Natta Catalyst," Macromolecules, 1992, 25, 2820 - 2827, it was confirmed that a soluble vanadium based Ziegler-Natta catalytic system  $\text{VOCl}_3/\text{Al}_2(\text{C}_2\text{H}_5)_3\text{Cl}_3$ , acts essentially as a single site catalyst although  $\text{VOCl}_3$  is not a metallocene. Homogeneous copolymers produced by such a catalyst system have been commercially available for several years. An example of such are the resins sold under the tradename Tafmer(TM) by Mitsui.

U.S. Patent No. 4,501,634 to Yoshimura et al is directed to an oriented, multilayered film which includes a Tafmer as a blend component in at least one layer.

Japanese Kokoku 37907/83 to Gunze Limited was directed to a heat-sealable biaxially oriented composite film wherein the heat seal layer contains Tafmer in a blend.

The foregoing patents disclose homogeneous ethylene alpha-olefins having densities below 0.90 g/cc. Heretofore, such resins have been generally unavailable in densities at and above 0.90 g/cc because of limitations in the  $\text{VOCl}_3$  process. Yet, U.S. Patent 1,209,825 to DuPont discloses homogeneous copolymers having densities up to about 0.920 made with single site Ziegler catalysts. When extruded into a film the 5 resins exhibit improved physical and optical properties as compared to films of heterogeneous copolymers. However, such homogeneous copolymers of densities above 0.90 g/cc have been commercially unavailable. It is believed that process limitations precluded the manufacture of such resins in any but bench-top 10 quantities. Metallocene catalysts, however, can provide such homogeneous copolymers in a wide range of densities in commercial quantities. Thus, the benefits of homogeneity can now be incorporated into copolymers having densities analogous to those of conventional VLDPEs and LLDPEs.

However, while providing improved physical properties such as optics, low extractables and improved impact, the narrow compositional distribution of some typical metallocene catalyzed resins can cause some processing difficulties. It has been found that such processing problems are avoided if some limited long 15 chain branching is introduced. That is, a typical metallocene catalyzed ethylene alpha-olefin may be thought of as a collection of linear chains, each of substantially identical length, each having approximately the same number of short chain (comonomer) branches distributed at regular intervals along that length. Splicing an abbreviated linear chain with the same regular comonomer distribution onto each of the linear 20 chains, or at least some of the chains in the collection, yields an ethylene alpha-olefin with essentially all of the physical properties of the original copolymer, but with an improved "body" or melt strength for improved processability including improved extrudability, orientation speeds and susceptibility to irradiation.

One way of quantifying the improved processability of the new homogeneous branched ethylene alpha-olefin copolymers is by the melt flow ratio  $l_{10}/l_2$  as described in ASTM D-1238. In WO 93/08221 Dow discloses such copolymers and notes that the  $l_{10}/l_2$  ratio for such is indicative of the degree of long chain branching. For linear polyolefins the  $l_{10}/l_2$  ratio increases as molecular weight distribution increases. 25 Molecular weight distribution may be defined as the ratio of weight average molecular weight (Mw) to number average molecular weight (Mn) or  $Mw/Mn$ . Conventional heterogeneous linear ethylene alpha-olefins typically have a high  $l_{10}/l_2$  and a correspondingly high molecular weight distribution. Linear homogeneous ethylene alpha-olefins have a low molecular weight distribution and a correspondingly low  $l_{10}/l_2$ . But 30 homogeneous ethylene alpha-olefins with long chain branching such as those supplied by Dow under the Insite trade name have low molecular weight distributions but high  $l_{10}/l_2$  ratios. In part, Dow defines these copolymers as having  $l_{10}/l_2 \geq 5.63$  and  $Mw/Mn \leq (l_{10}/l_2) - 4.63$ . This high  $l_{10}/l_2$  is an indication of ease of 35 processability discussed above, while the homogeneity, of which  $Mw/Mn$  is one measure, provides for improved physical properties also discussed above. A more detailed description of the chemistry and particularly the rheology of these branched homogeneous copolymers is given in WO 93/08221 which is incorporated herein by reference.

Dow in EP 416,815 disclosed the preparation of ethylene/olefin copolymers using monocyclopentadienylsilane complexed to a transition metal. The homogeneous ethylene copolymers which may be prepared using this catalyst are said to have better optical properties than typical ethylene polymers and be well suited for film or injection molding.

40 As will be shown below, it has been found that resins produced by the Dow process exhibit improved physical properties characteristic of single site catalyzed resins but also possess a processability similar to that of conventional Ziegler-Natta copolymers. It is believed that the Dow metallocene resins possess the limited long chain branching discussed above.

45 It is thus an object of the present invention to provide a film structure having improved physical properties such as improved optics and impact strength and low extractables.

It is a further object of the present invention to provide a film which is readily extruded and processed.

### Summary of the Invention

50 These as well as other objects are achieved by providing a film which includes a homogeneous single site catalyzed copolymer of ethylene and an alpha-olefin having from three to ten carbon atoms, wherein the single site catalyzed copolymer includes some long chain branching.

### Detailed Description of the Preferred Embodiments

55 The present invention is directed to a heat-shrinkable, thermoplastic film or bag containing a homogeneous copolymer of ethylene and at least one other alpha-olefin wherein the copolymer contains at least some limited long chain branching. The term "copolymer" as used herein is intended to denote polymers of

two or more comonomers. Therefore, although the present specification generally discusses ethylene alpha-olefin copolymers such term is intended to encompass copolymers of ethylene with one or more alpha-olefins or ethylene with an alpha-olefin and another comonomer.

The term "oriented" is used herein interchangeably with the term "heat shrinkable," these terms designating a material which has been stretched and set by cooling while substantially retaining its stretched dimensions. An oriented (i.e. heat shrinkable) material will tend to return to its original unstretched (unextended) dimensions when heated to an appropriate elevated temperature.

The film of the present invention is preferably oriented and is formed by extrusion processes especially art-known coextrusion methods. It is initially cooled to a solid state by, for example, cascading water or chilled air quenching, after which it is reheated to within its orientation temperature range and oriented by stretching. The stretching to orient may be accomplished in many ways such as, for example, by "blown bubble" techniques or "tenter framing." These processes are well known to those skilled in the art and refer to orientation procedures whereby the material is heated to its softening temperature and then stretched in the cross or transverse direction (TD) and/or in the longitudinal or machine direction (MD). After being heated and stretched, the film is quickly quenched while substantially retaining its stretched dimensions to cool the film rapidly and thus set or lock in the oriented molecular configuration.

The film layers may be formed by coextrusion, with additional layers thereafter being extrusion coated thereon to form multilayer films. Two multilayer tubes may also be formed with one of the tubes thereafter being extrusion coated or laminated onto the other. The extrusion coating method of film formation is preferable to coextruding the entire film when it is desired to subject one or more layers of the film to a treatment which may be harmful to one or more of the other layers. This may be done where it is desired to irradiate one or more layers of a film with high energy electrons where the film contains a barrier layer of one or more copolymers of vinylidene chloride (e.g., Saran(TM)), such as vinylidene chloride and vinyl chloride or vinylidene chloride and methyl acrylate as well as vinylidene chloride with ethyl acrylate or acrylonitrile.

Films of this type would, for example, comprise those where the barrier layer is a Saran(TM) layer in addition to or instead of an EVOH layer. Those skilled in the art generally recognize that irradiation with high energy electrons is generally harmful to such Saran(TM) barrier layer compositions, as irradiation may degrade and discolor Saran(TM), making it turn brown. Thus, if full coextrusion followed by high energy electron irradiation of the multilayer structure is carried out on a film having a barrier layer containing a Saran(TM) layer, the irradiation should be conducted at low levels and with care. Alternatively, this may be avoided by extruding a first layer or layers, subjecting the layer or layers to high energy electron irradiation and thereafter applying the Saran(TM) barrier layer and, for that matter, other layers (which may or may not have been irradiated) sequentially onto the outer surface of the extruded, previously irradiated, tube. This sequence allows for high energy electron irradiation of the first layer or layers without subjecting the Saran(TM) barrier layer to harmful discoloration.

Thus, as used herein the term "extrusion" or the term "extruding" is intended to include coextrusion, extrusion coating, or combinations thereof.

As noted, the present heat shrinkable film may optionally be subjected to an energetic radiation treatment, including, but not limited to corona discharge, plasma, flame, ultraviolet, and high energy electron treatment. Irradiation is most preferably performed prior to orientation and, in the case of a Saran-containing barrier structure, prior to extrusion coating of the barrier component. Radiation dosages are referred to herein in terms of the radiation unit "RAD," with one million RADS or a megarad being designated as "MR." A suitable radiation dosage of high energy electrons is in the range of up to about 12 MR, more preferably about 2 to about 9 MR. However, irradiation after orientation, regardless of structure, and performed at lower dosage levels, is also within the scope of the present invention.

For those embodiments in which the present heat shrinkable film is a multilayer film, each layer will generally serve some function or provide some characteristic to the overall structure. Seal layer composition will be chosen for ease of heat sealing and, depending on the intended end use application, other factors such as grease resistance may be of importance. Outer layer composition may be chosen for abuse resistance or, where a given end use application requires a folded over "lap" seal, sealability to the seal layer. If a barrier layer is required it will be chosen for the degree of gas or moisture impermeability needed for the ultimate product to be packaged. Further internal layers may serve to add bulk to the film, promote shrinkability, promote interlayer adhesion or any combination of these properties.

For purposes of the present invention it has been found that the use of homogeneous ethylene alpha-olefin copolymers with long chain branching in at least one core layer of a multilayer heat-shrinkable film provides improved impact properties and excellent shrink properties. More particularly, for a gas barrier material which includes a core layer of a polymeric material having gas barrier properties such as a

vinylidene chloride copolymer or an ethylene vinyl alcohol copolymer, it has been found that further internal layers of such homogeneous, branched ethylene alpha-olefin copolymers sandwiching the barrier layer provide for a machinable barrier film with improved impact resistance and free shrink. That is, it is within the scope of the present invention to provide a multilayer heat-shrinkable film having the general structure:

5      seal/core/barrier/core/abuse

wherein the respective compositions of the seal, barrier and abuse layers may be chosen from those polymeric resins or resin blends which provide the requisite functional properties. Examples of materials appropriate for use in the seal layer include EVA, LLDPE, VLDPE, EAA, EMAA, ionomers, homogeneous linear ethylene alpha-olefins and homogeneous branched ethylene alpha-olefins. Examples of materials

10     appropriate for use in the abuse layer include EVA, LLDPE, VLDPE, homogeneous linear ethylene alpha-olefins, homogeneous branched ethylene alpha-olefins, polypropylene, nylon, and high density polyethylene. Examples of materials appropriate for use in the barrier layer include vinylidene chloride-vinyl chloride, vinylidene chloride methyl acrylate, ethylene vinyl alcohol, acrylonitrile and nylon. However, other materials which have the needed properties of sealability and seal integrity, abuse resistance and low gas transmission

15     are also appropriate for use in the seal layer, abuse layer and barrier layer, respectively, of a film in accordance with the present invention. Each of the two core layers contain a homogeneous ethylene alpha-olefin copolymer either alone or in a blend with another polymeric material.

For those applications in which a homogeneous branched ethylene alpha-olefin copolymer provides adequate seal and abuse properties a film having the general structure:

20     seal/barrier/abuse

is also within the scope of the present invention wherein the seal and abuse layer each contain a homogeneous branched ethylene alpha-olefin either alone or in a blend with another polymeric material.

For those applications in which a barrier layer is not required, a film having the general structure:

seal/core/abuse

25     is also within the scope of the present invention wherein the core contains a homogeneous branched ethylene alpha-olefin either alone or in a blend. Further core layers of branched homogeneous ethylene alpha-olefins or other polymeric materials may also be included.

For each of the general structures set forth above it should be noted that further internal layers may be provided to promote inner layer adhesion or add bulk as may be necessary.

30     It has been found that a preferred method for producing a film in accordance with the present invention is an extrusion coating method such as described above. By such method a portion of the film structure is extruded, cooled and subjected to irradiation prior to be extrusion coated with the remaining layers and subsequently oriented. Preferably, the seal layer and a first core layer are extruded and irradiated prior to extrusion coating of the barrier layer, second core layer and abuse layer and subsequent orientation. Most 35     preferably the seal layer, first core layer and a third layer chosen to promote adhesion to the barrier layer are extruded and irradiated, followed by extrusion coating of the barrier layer, an adhesion promoting tie layer, the second core layer and the abuse layer and then orientation. For the present preferred method it has been found that the use of homogeneous branched ethylene alpha-olefins in the core layers reduces extrusion head pressure and allow for increased orientation speeds.

40     The following examples are representative of the preferred embodiments of the present films containing homogeneous branched ethylene alpha-olefin copolymers. In order to evaluate such films the following tests were employed:

Tensile Strength: A measure of the force required under constant elongation to break a specimen of the film; measured by ASTM D 882.

45     Elongation: A measure of the percent extension required to break a specimen of the film; measured by ASTM D 882.

Modulus: The ratio of the change in force to the change in elongation in the straight line portion of an Instron Tensile Testing curve; measured by ASTM D 882 - Method A.

50     Tear Propagation: The force required to propagate a tear from a tiny slit made by a sharp blade in a specimen of the film; measured by ASTM D 1938.

Free Shrink: The percent dimensional change in a 10 cm. x 10 cm. specimen of film when subjected to a selected heat; measured by ASTM D 2732.

Ball Burst: The energy necessary to burst and penetrate a restrained specimen of film; measured by ASTM D 3420.

55     Instrumented Impact: The energy necessary to puncture a restrained specimen of film, similar to ball burst, defined above. However, the Instrumented Impact Tester has the ability to measure the tensile/elongation curve to break. The "gradient" is the ratio of the change in force to change in elongation in the straight line portion of the curve. "Peak" is a measure of the maximum force exerted on the specimen to impart rupture.

"Impact Energy" is a measure of the energy absorbed by the sample prior to rupture. Instrumented impact is measured by ASTM D 3763.

Haze: The percentage of transmitted light which is scattered forward while passing through a specimen; measured by ASTM D 1003 - Method A.

5 Clarity: A measure of the distortion of an image viewed through a specimen; measured by ASTM D 1746.

Gloss: The surface reflectance or shine of a specimen; measured by ASTM D 2457.

Parallel Plate: A bag is confined between two plates a specified distance apart and is inflated until its seal fails. The pressure level inside the bag at the point of failure is a measure of seal quality. Results are reported in inches of water pressure (IOWP).

10 LRHB (Linear Ramped Hot Burst): A clean sealed bag is inflated to a specified dwell pressure and the seal area is submerged in hot water at 182 °F. After five seconds the pressure inside the bag is increased at the rate of 2 inches of water/second. The time to failure and burst pressure is a measure of seal quality. Test results are reported in seconds and inches of water pressure (IOWP).

15 LRHB-G (Linear Ramped Hot Burst - Grease): The procedure is the same as the LRHB test described above except peanut oil is first applied to the seal area.

VPHB (Variable Pressure Hot Burst): As with the LRHB test described above, a clean sealed bag is inflated to a specified dwell pressure and the seal area is submerged in hot water at 182 F. After five seconds the pressure inside the bag is increased at a specified rate ranging 1 to 7 inches of water/second. Here again, test results are reported as seconds and inches of water pressure (IOWP).

20 VPHB-G (Variable Pressure Hot Burst - Grease) : The procedure is the same as the VPHB test described above except peanut oil is first applied to the seal area.

25 Gel: A measure of the relative amount of ultra high molecular weight polymer present in a sample. A gel measurement can provide an indication of the level of crosslinking which is present in a sample since the amount of polymer which is collected as gel increases with crosslinking. Gel is determined by solvent extraction with boiling toluene. In this method a specimen weighting 0.4 grams is extracted for 21 hours in a cellulose thimble, removed, dried and reweighed. The percentage of gel is calculated by ratioing the polymer remaining (toluene insoluble fraction) to the original weight. However, a gel value of 0% cannot reliably indicate that no crosslinking has occurred. Rather, the level of crosslinking may not be great enough to provide measurable gel.

30 DSC: The differential scanning calorimeter (DSC) is an instrument which measures the heat flow to a polymer sample during the programmed heating of it at 10 °C per minute. The trace obtained from the DSC can be used to characterize a sample's starting and ending melting point as well as its peak melting point(s).

35 Polydispersity (Mw/Mn): A measure of the uniformity of chain lengths within a polymer sample. It defines the breadth of molecular weight distribution. It is obtained by dividing the weight average molecular weight (Mw) by the number average molecular weight (Mn). The Mw and Mn are determined by Gel Permeation Liquid Chromatography.

#### Example 1

40 A two layer coextruded precursor film having the structure inner layer/outermost layer was formed. The inner layer was a blend of 90% by weight of NA 295-000, a 6.7% vinyl acetate EVA supplied by Quantum, and 10% by weight of Dowlex 2045. The outermost layer was a blend of 85% by weight of XUR-1567-48562-B9, a homogeneous ethylene octene copolymer having a density of 0.904 g/cc and a 1.0 MI with 45 limited long chain branching sold on a developmental basis by DOW, and 15% by weight of EA 719-009, an ethylene butyl-acrylate copolymer having 18.5% by weight butyl-acrylate supplied by Quantum. Following irradiation, the precursor film was extrusion coated with a barrier layer of a vinylidene chloride methyl acrylate from Dow and an outer abuse layer of a blend of 92.5% by weight of LD 318.92 from Exxon, a 9% vinyl acetate EVA and 7.5% by weight of Dowlex 2045.

50 The resultant four layer film was then oriented by a trapped bubble method out of hot water.

#### Example 2

55 The procedure set forth in Example 1 was repeated with the exception that the outermost layer of the precursor film was 85% by weight of XUR-1567-48562-B9 from Dow and 15% by weight of Nucrel 1202HC, an ethylene methacrylic acid supplied by DuPont.

Example 3

The procedure set forth in Example 1 was repeated with the exception that the outermost layer of the precursor film was 85% by weight of XUR-1567-48562-B9 from Dow and 15% by weight of Bynel CXA 5101, an anhydride grafted EVA having 18.4% vinyl acetate supplied by DuPont.

Example 4

The procedure set forth in Example 1 was repeated with the exception that the outermost layer of the precursor film was 85% by weight of XUR-1567-48562-B9 from Dow and 15% by weight of LD 318.92 from Exxon, a 9% vinyl acetate EVA.

Example 5

For comparative purposes, the procedure set forth in Example 1 was repeated with the exception that the outermost layer of the precursor film was 85% by weight of Dowlex 2045 and 15% by weight of EA 719-009 an 18.5% butyl acrylate EBA.

Examples 6-10

The oriented multilayer films of Examples 1-5 were tested for instrumented impact, free shrink and "peel", a measure of bond strength between the outermost layer of the precursor film and the barrier layer which has been extrusion coated onto it. The results are given below in Table I. The instrumented impact and T peel numbers for the films of Examples 1-4 compare favorably to those of comparative Example 5. The free shrink of the films embodying the present invention are superior to those of the prior art structure. It should be noted that the numbers given for comparative Example 5 are representative for a series of production runs.

Table I

Ex. #	Film of	Instrumented Impact		Free Shrink L & T	T Peel lbs/in
		Peak(lbs)	Energy(ft)		
6	1	51.0	2.35	73	0.143
7	2	55.7	2.62	73	0.157
8	3	49.3	2.32	75	0.214
9	4	54.0	2.70	71	0.133
10	5	51.3	2.20	49.9	0.180

Example 11-19

The films of Examples 11-19 each are formed of a precursor film having an inner layer of 90% NA 295-000, a 6.7% EVA supplied by Quantum and 10% Dowlex 2045, a core layer and an outermost adhesive layer of EP 4062-3, a 15% vinyl acetate EVA supplied by DuPont. Following irradiation at a selected dosage the precursor film is extrusion coated with a barrier layer of a vinylidene chloride methyl acrylate from Dow and an outer abuse layer of 92.5% LD 318.92 and 7.5% Dowlex 2045.

Thereafter the total structure was oriented out of hot water by a trapped bubble technique with constant orientation preheat and hot bath temperatures of 195 °F.

The core layer composition and irradiation dosage for each structure are given below in Table II. Also shown are orientation speeds. It should be noted that for the films of Examples 11-16, orientation speed increases with increasing irradiation dosage, whereas such relationship is not seen for comparative Examples 17-19.

The resins employed in the core layer are XUR-1567-48562-B9, a homogeneous ethylene octene copolymer having a density of 0.904 g/cc and a 1.0 MI with limited long chain branching supplied by Dow represented by B9 (below), XUR-1567-48562-B4, a homogeneous ethylene octene copolymer having a density of 0.909 g/cc and a 1.0 MI with limited long chain branching supplied by Dow represented by B4

The film structure and target percent thickness per layer for each film was:

5  
10  
15  
20  
25  
30  
35  
40  
45  
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11.9	49.6	6.0	8.7	23.8
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Inner/Core/Adhesive//Barrier/Abuse

Table II

Ex. #	Core	Dosage MR	Orientation Speed feet/min.
11	89	2	46
12	89	4	53
13	89	6	53
14	84	2	40
15	84	4	46
16	84	6	51
17	4203	2	46
18	4203	4	46
19	4203	6	46

Examples 20-28

The oriented multilayer films of Examples 11-19 were tested for instrumented impact and free shrink. The results are given below in Table III. The films of Examples 11-16 compare favorably in all values to those of comparative Examples 17-19.

Table III

Ex.	Film of	Instrumented Impact		Free Shrink L & T
		Peak(lbs)	Energy(ft.lbs)	
20	11	59.6	3.45	72
21	12	59.9	3.08	71
22	13	61.5	3.17	74
23	14	63.0	3.36	68
24	15	60.4	2.96	67
25	16	64.6	3.39	69
26	17	52.8	2.83	76
27	18	55.3	2.87	74
28	19	56.7	3.01	77

EXAMPLE 29

A three layer coextruded precursor film having the structure: inner layer/core layer/outermost layer was formed. The inner layer was a blend of 90% by weight of NA 295-000, a 6.7% vinyl acetate EVA supplied by Quantum, 10% by weight of Dowlex 2045, a 0.920 density heterogeneous linear ethylene octene copolymer from Dow. The outermost layer was EP 4062-2, a 15% vinyl acetate EVA supplied by DuPont. The core layer was XU59220.01, lot 427833 a homogeneous ethylene octene copolymer having a density of 0.9016 g/cc and a 0.9 MI with long chain branching sold on a developmental basis by Dow. The resin has an  $I_{10/2}$  of 10.9 and a  $M_w/M_n$  of 2.03.

Following irradiation, the precursor film is extrusion coated with a carrier layer of a vinylidene chloride-methyl acrylate from Dow blended with an epoxidized soybean oil and an ethylene methyl acrylate, an adhesive layer of Elvax 3175GC, a 28% vinyl acetate EVA from DuPont, a core layer of XU 59220.01, lot 427833 from Dow and an outer abuse layer of a blend of 92.5% by weight of LD 318.92 from Exxon, a 9% vinyl acetate EVA, and 7.5% by weight of Dowlex 2045.

The resultant seven layer film was then oriented by a trapped bubble method out of hot water. The preheat temperature 192°F and the hot bath was 197°F. The final oriented structure was 1.8 mil thick with the individual layers having target thicknesses as follows:

10 seal/core/outer//barrier/tie / core/abuse  
0.28 0.75 0.13 0.18 0.13 0.18 0.15

15 EXAMPLE 30

The procedure set forth in Example 29 was repeated with a final oriented structure having a thickness of 2.2 mils. The individual layers had a target thickness as follows:

20 seal/core/outer//barrier/tie / core/abuse  
0.33 1.0 0.14 0.18 0.14 0.24 0.17

25 EXAMPLE 31

The procedure set forth in Example 29 was repeated with a final oriented structure having a thickness of 2.7 mils. The individual layers had a target thickness as follows:

30 seal/core/outer//barrier/tie / core/abuse  
0.36 1.39 0.15 0.18 0.15 0.30 0.17

35 EXAMPLE 32

40 The procedure set forth in Example 29 was repeated with the following exceptions. Preheat temperature was 191°F and the hot bath was 184°. The final oriented structure had a thickness of 3.0 mils. The individual layers had a target thickness as follows:

45 seal/core/outer//barrier/tie/core/abuse  
0.36 1.59 0.15 0.18 0.15 0.39 0.17

50 EXAMPLE 33

The procedure of Example 32 was repeated with the only exception being a difference in orientation condition. The preheat temperature was 199°F and the hot bath was 195°F.

55 EXAMPLE 34

The procedure set forth in Example 29 was repeated with the core layers being replaced with XU59243.00, a homogeneous ethylene octene copolymer having a density of 0.904 g/cc and a 0.8 MI with

long chain branching sold on a developmental basis by Dow. The resin has an  $I_{10}/I_2$  of 9.2 and Mw/Mn of 2.17. For orientation, the preheat temperature was 194°F and the hot bath was 190°F. The final oriented structure had a thickness of 3.0 mils. The individual layers had target thicknesses as follows:

5

seal/core/outer//barrier/tie / core /abuse  
.36 1.59 .15 .18 .15 .39 .17

10

EXAMPLE 35

15 The procedure of Example 34 was repeated with the only exception being different orientation conditions. The preheat temperature was 200° and the hot bath was 195°F.

EXAMPLE 36

20 The procedure set forth in Example 29 was repeated with the core layers being replaced with XU 59220.00, a homogeneous ethylene octene copolymer having a density of 0.904 g/cc and a 0.9 MI with long chain branching sold on a developmental basis by Dow. The resin has an  $I_{10}/I_2$  of 11.3 and an Mw/Mn of 2.4. The preheat temperature was 194°F and the hot bath was 190°F. The final oriented structure had a thickness of 3.0 mils. The individual layers had target thicknesses as set forth in Example 32 above.

25 EXAMPLE 37

The procedure of Example 36 was repeated with the only exception being a difference in orientation conditions. The preheat temperature was 200°F and the hot bath was 195°F.

30 EXAMPLE 38

35 The procedure set forth in Example 29 was repeated with the core layers being replaced with XU 59220.01, lot 421733, a homogeneous ethylene octene copolymer having a density of 0.9028 and a 0.9 MI with long chain branching sold on a developmental basis by Dow. The resin has an  $I_{10}/I_2$  of 10.5 and an Mw/Mn of 2.4. The preheat temperature was 194°F and the hot bath was 190°F. The final oriented structure had a thickness of 3.0 mils and individual layers as set forth in Example 32 above.

EXAMPLE 39

40 The procedure of Example 38 was repeated with the only exception being a difference in orientation conditions. The preheat temperature was 200°F and the hot bath was 195°F.

EXAMPLE 40

45 The procedure set forth in Example 29 was repeated with the core layers being replaced with XU 59220.02, a homogeneous ethylene octene copolymer having a density of 0.906 and a 0.8 MI with long chain branching sold on a developmental basis by Dow. The resin has an  $I_{10}/I_2$  of 11.8 and an Mw/Mn of 2.2. The preheat temperature was 197°F and the hot bath was 194°F. The final oriented structure and individual layers had its thickness of 3.0 mils and individual layer as set forth in Example 32 above.

50

EXAMPLE 41

The procedure of Example 40 was repeated with the only exception being a difference in orientation condition. Preheat temperature was 200°F and the hot bath was 195°F.

55

EXAMPLE 42

5 The procedure set forth in Example 29 was repeated with the core layers being replaced with XU 59220.03, a homogeneous ethylene octene copolymer having a density of 0.898 and a 0.9 MI with long chain branching sold on a developmental basis by Dow. The resin has an  $I_{10}/I_2$  of 10.5 and an Mw/Mn of 2.0. The preheat temperature was 187°F and the hot bath was 181°F. The final structure and individual layers had thicknesses as set forth above in Example 32.

EXAMPLE 43

10 The procedure of Example 42 with the only exception being a difference in orientation condition. The preheat temperature was 200°F and the hot bath was 195°F.

EXAMPLE 44

15 The procedure set forth in Example 29 was repeated with the core layers being replaced with XU 59220.04, a homogeneous ethylene octene copolymer having a density of .896 and a 0.9 MI with long chain branching sold on a developmental basis by Dow. The resin has an  $I_{10}/I_2$  of 10.5 and an Mw/Mn of 2.17. The preheat temperature was 188°F and the hot bath was 182°F. The final oriented structure and 20 individual layers had thicknesses as set forth in Example 32 above.

EXAMPLE 45

25 The procedure of Example 44 was repeated with the only exception being a difference in orientation condition. The preheat temperature 200°F and the hot bath was 195°F.

EXAMPLE 46

30 The procedure set forth in Example 29 was repeated with the core layers being replaced with Insite C24, a homogeneous branched ethylene octene copolymer having a density of 0.904 g/cc and a 0.9 MI sold on a developmental basis by Dow. The resin has an  $I_{10}/I_2$  of 9.2 and an Mw/Mn of 2.0. The preheat temperature was 200°F and the hot bath was 195°F. The final structure was 3.0 mils and individual layers had target thicknesses as set forth in Example 32 above.

35 EXAMPLE 47

For comparative purposes, the procedure set forth in Example 29 was repeated with the core layers being replaced with Attane 4203, a heterogeneous octene copolymer from Dow having a density of 0.905 g/cc and a 0.8 MI. The resin has an  $I_{10}/I_2$  of 8.2 and an Mw/Mn of 3.8. The preheat temperature was 194°F 40 and the hot bath was 190°F. The final oriented structure had a thickness of 3.0 mils and individual layers had target thicknesses as set forth above in Example 32.

EXAMPLE 48

45 For comparative purposes, the procedure of Example 47 was repeated with the only exception being a difference in orientation condition. The preheat temperature was 200°F and hot bath was 195°F.

EXAMPLES 49 - 68

50 To evaluate their physical properties, the oriented multilayer films of Examples 29-48 were tested for instrumented impact and free shrink. Examples 34 and 35 are based on core layers of XU 59243.00, a homogeneous branched ethylene octene from Dow which has a density of 0.904 g/cc and a 0.8 MI, a density and melt index essentially the same as that for Attane 4203, the heterogeneous linear ethylene octene copolymer employed in Comparative Examples 47 and 48. However, the film structures of Examples 55 34 and 35 show greatly improved peak load and energy to break as compared to the films of Examples 47 and 48, respectively. The results are given in Table IV below.

EX.# L&T	FILM OF	PEAK LOAD (lbs)	ENERGY TO BREAK (ft. lbs)	FREE SHRINK
5	49	29	2.3	83
	50	30	2.9	80
	51	31	4.2	81
	52	32	6.2	86
	53	33	4.9	76
10	54	34	9.3	75
	55	35	8.0	67
	56	36	7.4	76
	57	37	5.4	70
	58	38	5.1	77
15	59	39	6.0	73
	60	40	5.4	68
	61	41	6.0	66
	62	42	6.5	96
	63	43	5.0	78
20	64	44	6.3	96
	65	45	4.9	80
	66	46	10.6	77
	67	47	5.1	79
	68	48	5.6	74

25

EXAMPLES 69-88

30 To evaluate the processability of the homogeneous branched ethylene octene copolymers found in each of the above film structures, the orientation speed was noted for each structure at each orientation temperature. Surprisingly, the film structures containing the homogeneous branched ethylene octene copolymers from Dow which showed improved toughness in Examples 49 - 66 above generally oriented faster than the films of Comparative Examples 47 and 48. The results are given in Table V below.

35

40

45

50

55

Table V

EX.#	FILM OF	PREHEAT (°F)	HOT BATH (°F)	ORIENTATION SPEED (ft/min)
69	29	192	197	51
70	30	192	197	53
71	31	192	197	50
72	32	191	184	32
73	33	199	195	45
74	34	194	190	33
75	35	200	195	40
76	36	194	190	30
77	37	200	195	37
78	38	194	190	33
79	39	200	195	40
80	40	197	194	32
81	41	200	195	35
82	42	187	181	40
83	43	200	195	51
84	44	188	182	36
85	45	200	195	46
86	46	200	195	53
87	47	194	190	32
88	48	200	195	36

EXAMPLES 89-90

To evaluate the processability of the homogeneous long chain branched ethylene octene copolymers incorporated into the present film structures, the head pressures were noted during extrusion of both the substrate and the coating of the films of Example 34 and Comparative Example 47. Examples 29-33 and 36-45 are not included in Table VI below because valid comparisons can only be made between resins of the same melt index. Example 46 is not included because it was not made under comparable extrusion conditions. It should be noted that although the films of Examples 35 and 48 contain the same polymeric materials as those of Examples 34 and 47, evaluated below, they are not included because they result from the same extrusion runs. That is, Example 35 is the orientation of the unoriented tape structure described in Example 34 under differing orientation conditions from those described in Example 34. The same can be said for Example 48 with respect to Example 47. The homogeneous branched ethylene octene copolymer employed in Example 34 provided lower extrusion head pressures than the heterogeneous linear ethylene octene copolymer of Example 47.

Table VI

EX.#	FILM OF	HEAD PRESSURE OF 1ST EXTRUDER (psi)	HEAD PRESSURE OF 2ND EXTRUDER (psi)
89	34	5230	5750
90	47	5600	6267

EXAMPLE 91

The procedure set forth in Example 29 was repeated with the two core layers being replaced with a blend of 80% by weight of XU59243.00, a homogeneous ethylene octene copolymer having a density of 0.904 g/cc and a 0.8 MI with long chain branching sold on a developmental basis by Dow, and 20% by weight of LD-318.92, a 9% VA ethylene vinyl acetate sold by Exxon. The XU59243.00 resin has an  $I_{10}/I_2$  of

5 The final oriented structure had a thickness of 3.0 mils. The individual layers had a target thickness as follows:

10 Seal/Core/Outer//Barrier/tie/Core/Abuse  
0.36 1.59 0.15 0.18 0.15 0.39 0.17

15

EXAMPLE 92

15 The procedure set forth in Example 29 was repeated with the two core layers being replaced with a blend of 90% by weight of XU 59243.00 and 10% by weight of LD-318.92 from Exxon.

16 The final oriented structure had a thickness of 3.0 mils. The individual layers had a target thickness as follows:

20 Seal/Core/Outer//Barrier/tie/Core/Abuse  
0.36 1.59 0.15 0.18 0.15 0.39 0.17

25 EXAMPLE 93

25 The procedure set forth in Example 29 was repeated with the two core layers being replaced with a blend of 80% by weight of XU59243.00 and 20% by weight of Elvax 3175GC, a 28% VA ethylene vinyl acetate sold by DuPont

30 The final oriented structure had a thickness of 3.0 mils. The individual layers had a target thickness as follows:

35 Seal/Core/Outer//Barrier/tie/Core/Abuse  
0.36 1.59 0.15 0.18 0.15 0.39 0.17

40 EXAMPLE 94

40 The procedure set forth in Example 29 was repeated with the two core layers being replaced with a blend of 90% by weight of XU59243.00 and 10% by weight of Elvax 3175GC from DuPont.

45 The final oriented structure had a thickness of 3.0 mils. The individual layers had a target thickness as follows:

50

Seal/Core/Outer//Barrier/tie/Core/Abuse  
0.36 1.59 0.15 0.18 0.15 0.39 0.17

EXAMPLE 95

55 The procedure set forth in Example 29 was repeated with the two core layers being replaced with a blend of 85% by weight of XU59243.00 and 15% by weight of Escorene LD-720.92, a 19% VA ethylene vinyl acetate sold by Exxon.

follows:

5 Inner/Core/Outer//Barrier/tie/Core/Abuse  
0.36 1.59 0.15 0.18 0.15 0.39 0.17

**10 EXAMPLE 96-100**

The oriented multilayer films of Examples 91-95 were tested for instrumented impact and free shrink. The results are given below in Table VII.

15

Table VI

EX.#	FILM OF	INSTRUMENTED IMPACT		FREE SHRINK L&T	
		PEAK LOAD (lbs)	ENERGY TO BREAK (ft. lbs.)		
20	96	91	84	5.6	74
	97	92	86	5.8	76
25	98	93	84	5.0	82
	99	94	96	6.2	80
	100	95	85	5.1	80

The foregoing description of preferred embodiments of the invention have been presented for purposes of illustration and description. It is not intended to be exhaustive or to limit the invention to the precise form disclosed, and modifications and variations are possible in light of the above teachings or may be acquired 30 from practice of the invention. The embodiment were chosen and described in order to explain the principles of the invention and its practical application to enable one skilled in the art to utilize the invention in various embodiments and with various modifications as are suited to the particular use contemplated. It is intended that the scope of the invention be defined by the claims appended hereto, and their equivalents.

35 Claims

40 1. A film suitable for packaging comprising a homogeneous single site catalyzed copolymer of ethylene and an alpha-olefin having from three to ten carbon atoms, said single site catalyzed copolymer having limited long chain branching.

2. A film as set forth in claim 1 wherein said film is heat-shrinkable.

3. A heat shrinkable film as set forth in claim 1, wherein said alpha-olefin has from four to eight carbon atoms.

45 4. A film as set forth in claim 1 wherein said single site catalyzed copolymer is blended with another thermoplastic homopolymer or copolymer.

50 5. A film as set forth in claim 1 wherein said single site catalyzed copolymer has a density of from about 0.86 g/cc to about 0.95 g/cc.

6. A film as set forth in claim 1 wherein said film is a multilayer film.

55 7. A heat shrinkable multilayer film comprising at least one inner core layer comprising a homogeneous single site catalyzed copolymer of ethylene and an alpha-olefin having from four to ten carbon atoms and having at least some long chain branching said copolymer having a density of from about 0.89 g/cc to about 0.91 g/cc.

octene.

9. The heat shrinkable film of claim 7 wherein said homogeneous branched copolymer is blended with another thermoplastic homopolymer or copolymer.  
5
10. The heat shrinkable film of claim 9 wherein said other thermoplastic homopolymer or copolymer is a copolymer of ethylene and a second comonomer selected from the group consisting of vinyl acetate, alkyl acrylate, carbon monoxide, butadiene, styrene, acrylic acid, methacrylic acid, a metal neutralized salt of an acrylic acid, and an alpha-olefin.  
10
11. A heat shrinkable multilayer film comprising at least two core layers, each of said core layers comprising a homogeneous single site catalyzed copolymer of ethylene and an alpha-olefin having from four to ten carbon atoms and having at least some long chain branching said copolymer having a density of from about 0.89 to about 0.91.  
15
12. The heat shrinkable film set forth in claim 11 wherein the homogeneous branched copolymer of one of said two core layers is identical to the homogeneous branched copolymer of the other of said layers.  
20
13. The heat shrinkable film set forth in claim 11 wherein the alpha-olefin having from four to ten carbon atoms is octene.  
25
14. The heat shrinkable film of claim 11 wherein said homogeneous branched copolymer is blended with another thermoplastic homopolymer or copolymer.  
30
15. The heat shrinkable film of claim 11 wherein said other thermoplastic homopolymer or copolymer is a copolymer of ethylene and a second comonomer selected from the group consisting of vinyl acetate, alkyl acrylate, carbon monoxide, butadiene, styrene, acrylic acid, methacrylic acid, a metal neutralized salt of an acrylic acid, and an alpha-olefin.  
35
16. The heat shrinkable film of claim 11 wherein the homogeneous branched copolymer has a density of from about 0.90 g/cc to about 0.91 g/cc.  
40
17. A heat shrinkable multilayer film having the general structure:  
45 seal/core/barrier/core/abuse  
wherein each of the core layers comprise the same homogeneous long chain branched single-site catalyzed copolymer of ethylene and an alpha-olefin having from four to ten carbon atoms, said copolymer having a density of from about 0.89 g/cc to about 0.91 g/cc.
18. The heat shrinkable film of claim 17 wherein said alpha-olefin having from four to ten carbon atoms is octene.  
50
19. The heat shrinkable film of claim 17 wherein said homogeneous branched copolymer is blended with another thermoplastic homopolymer or copolymer.  
55
20. The heat shrinkable film of claim 19 wherein said other thermoplastic homopolymer or copolymer is a copolymer of ethylene and a second comonomer selected from the group consisting of vinyl acetate, alkyl acrylate, carbon monoxide, butadiene, styrene, acrylic acid, methacrylic acid, a metal neutralized salt of an acrylic acid, and an alpha-olefin.  
60
21. The heat shrinkable film of claim 17 wherein the homogeneous branched copolymer has a density of from about 0.90 g/cc to about 0.91 g/cc.  
65
22. The heat shrinkable film of claim 17 wherein the barrier layer comprises a vinylidene chloride copolymer.  
70
23. The heat shrinkable film of claim 22 wherein the barrier layer comprises a vinylidene chloride-methyl acrylate copolymer.  
75

comonomer selected from the group consisting of vinyl acetate, alkyl acrylate, acrylic acid, methacrylic acid, a metal neutralized salt of an acrylic acid, and an alpha olefin.

5 25. The heat shrinkable film set forth in claim 17 further including additional internal layers to promote interlayer adhesion.

10 26. A heat shrinkable multilayer film comprising:  
a) a seal layer;  
b) a first core layer comprising a homogeneous, long chain branched ethylene alpha-olefin copolymer having a density of from about 0.89 g/cc to about 0.91 g/cc;  
c) a barrier layer;  
d) a second core layer comprising a homogeneous, long chain branched ethylene alpha-olefin copolymer having a density of from about 0.89 g/cc to about 0.91 g/cc; and  
e) an abuse layer;  
wherein the homogeneous, branched ethylene alpha-olefin of the first core layer differs from that of the second core layer.

15 27. The heat shrinkable film claim 26 further including additional internal layers to promote interlayer adhesion.

20 28. A multilayer heat shrinkable film comprising:  
a) a sealing layer comprising a homogeneous, single site catalyzed copolymer of ethylene and a alpha-olefin with four to ten carbon atoms, said copolymer having long chain branching and a density of from about 0.89 g/cc to about 0.91 g/cc;  
b) a barrier layer; and  
c) an abuse layer comprising a homogeneous single site catalyzed copolymer of ethylene and a alpha-olefin with four to ten carbon atoms, said copolymer having long chain branching and a density of from about 0.89 g/cc to about 0.91 g/cc.

25 30 29. A heat shrinkable multi-layer film having the general structure:  
seal/core/barrier/core/abuse  
when each of the two core layers comprise the same homogenous copolymer of ethylene and octene having an  $l_{10}/l_2$  greater than or equal to 5.63 and an  $Mw/Mn$  less than or equal to  $(l_{10}/l_2) - 4.63$ , said copolymer having a density of from about 0.89 g/cc to about 0.91 g/cc.

35 30. The heat shrinkable film of claim 29 wherein at least one of said core layers further includes another thermoplastic homopolymer or copolymer blended with the homogenous ethylene octene polymer.

40 31. The heat shrinkable film of claim 30 wherein said other thermoplastic homopolymer or copolymer is a copolymer of ethylene and a second comonomer selected from the group consisting of vinyl acetate, alkyl acrylate, carbon monoxide, butadiene, styrene, acrylic acid, methylacrylic acid, a metal neutralized salt of an acrylic acid and an alpha olefin.

45 32. The heat shrinkable film of claim 31 further including additional internal layers to promote interlayer adhesion.

50 33. The heat shrinkable multilayer film comprising:  
a) a sealing layer;  
b) a first core layer comprising a homogeneous ethylene octene copolymer having an  $l_{10}/l_2$  greater than or equal to 5.63 and an  $Mw/Mn$  less than or equal to  $(l_{10}/l_2) - 4.63$  and having a density of from about 0.89 g/cc to about 0.91 g/cc;  
c) a barrier layer;  
d) a second core layer comprising a homogenous ethylene octene copolymer having an  $l_{10}/l_2$  greater than or equal to 5.63 and an  $Mw/Mn$  less than or equal to  $(l_{10}/l_2) - 4.63$  and having a density of from about 0.89 g/cc to about 0.91 g/cc;  
e) an abuse layer.

copolymer of said first core layer differs from that of said second core layer.

35. The heat shrinkable film set forth in claim 33 further including additional internal layers to promote interlayer adhesion.  
5
36. The heat shrinkable film set forth in claim 33 wherein the homogenous ethylene octene copolymer of at least one of said first and second core layers is blended with another thermoplastic homopolymer or copolymer.  
10
37. The heat shrinkable film set forth in claim 36 wherein said other thermoplastic homopolymer or copolymer is a copolymer of ethylene and a second comonomer selected from the group consisting of vinyl acetate, alkyl acrylate, carbon monoxide, butadiene, styrene, acrylic acid, methacrylic acid, a metal neutralized salt of an acrylic acid and an alpha olefin.  
15
38. A heat shrinkable multilayer film having the general structure:  
seal/core/abuse  
wherein the core layer comprises a homogeneous long chain branched single site catalyzed copolymer of ethylene and an alpha-olefin having from four to ten carbon atoms, said copolymer having a density of from about 0.89 g/cc to about 0.91 g/cc.  
20
39. The heat shrinkable film of claim 38 further including a second core layer, said core layer comprising a homogeneous long chain branched single site catalyzed copolymer of ethylene and an alpha-olefin having from four to ten carbon atoms, said copolymer having a density of from about 0.89 g/cc to about 0.91 g/cc.  
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DOCUMENTS CONSIDERED TO BE RELEVANT			
Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.Cl.)
X	WO-A-92 14784 (EXXON CHEMICAL PATENTS INC.)	1,3-6, 28,29,33	832B27/32 C08J5/18
Y	* page 3, line 30 - page 10, line 30; claims 1,4,5,8,12,15,17 *	2,7-39	865D65/40
P,X	MODERN PLASTICS INTERNATIONAL vol. 23, no. 8 , August 1993 , LAUSANNE CH pages 40 - 41 DON SCHWANK 'Single-site metallocene catalysts yield tailor-made polyolefin resins'	1,3,5, 29,33	
X	EP-A-0 495 099 (MITSUI PETROCHEMICAL INDUSTRIES, LTD.)	1,3,5,6, 28,29,33	
Y	* page 3, line 15 - line 40; claims 1,7 * * page 6, line 20 - line 37 * * page 20, line 16 - line 57 *	2,7-27	
X	EP-A-0 416 815 (THE DOW CHEMICAL COMPANY)	1,3,5, 29,33	
	* page 11, line 32 - page 12, line 48; claim 31; figure 16; examples 12-32; table 2 *		
X	EP-A-0 452 920 (MITSUI PETROCHEMICAL IND., LTD.)	1,3,5	C08J C08F C08L 832B 865D
	* page 2, line 20 - line 54; claim 1 * * page 5, line 1 - page 6, line 40 * * page 14, line 37 - page 16, line 1 *		
Y	GB-A-2 206 890 (W R GRACE & CO - CONN)	2,7-39	
	* page 11, line 32 - page 14, line 22; claims 1,5,15,20 *		
Y	US-A-5 132 074 (ISOZAKI ET AL.)	2,7-39	
	* column 3, line 27 - column 4, line 15; claim 1; example 1 *		
		-/-	
The present search report has been drawn up for all claims			
Place of search	Date of completion of the search	Examiner	
THE HAGUE	17 February 1994	Derz, T	
CATEGORY OF CITED DOCUMENTS			
X : particularly relevant if taken alone	T : theory or principle underlying the invention		
Y : particularly relevant if combined with another document of the same category	E : earlier patent document, but published on, or after the filing date		
A : technological background	D : document cited in the application		
O : non-written disclosure	L : document cited for other reasons		
P : intermediate document	& : member of the same patent family, corresponding document		



DOCUMENTS CONSIDERED TO BE RELEVANT

Category	Citation of document with indication, where appropriate, of relevant passages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int.CLS)
P, X	WO-A-93 08221 (THE DOW CHEMICAL COMPANY) * page 4, line 2 - page 11, paragraph 1; claims 2,8,10,27-30; examples 5,20; tables 2,3 *	1-3,5, 29,33	
D, Y	US-A-5 059 481 (LUSTIG ET AL.) * the whole document *	17-33	

TECHNICAL FIELDS  
SEARCHED (Int.CLS)

The present search report has been drawn up for all claims

Place of search	Date of completion of the search	Examiner
THE HAGUE	17 February 1994	Derz, T
<b>CATEGORY OF CITED DOCUMENTS</b>		
X : particularly relevant if taken alone Y : particularly relevant if combined with another document of the same category A : technological background G : non-written disclosure P : intermediate document	T : theory or principle underlying the invention E : earlier patent document, but published on, or after the filing date D : document cited in the application L : document cited for other reasons	A : member of the same patent family, corresponding document